## REMARKS

Claims 17-73 were previously pending in the application. Claims 17-19, 22-69, and 72-73 are canceled. Claims 20-21 and 70-71 remain pending in the present application. No new matter is entered, and the amendments made herein are fully supported by the specification, claims and drawings as originally filed.

Claims 20, 48, 49, 70 and 71 are objected for reciting "%" rather than "wt. %" as per the specification. Claims 48 and 49 are cancelled. Claims 20, 70 and 71 are amended as suggested in the Office Action. Applicants request that the objection be withdrawn.

The Examiner rejected claims 17-19, 30-44, 47, 50-66, 69 and 72-73 under 35 U.S.C. § 103(a) as being unpatentable over U.S. Patent No. 6,607,620 to Knickerbocker et al. in view of U.S. Patent No. 5,130,067 to Flaitz et al. Applicants traverse the rejection to the extent that it can be maintained. As these claims have been canceled, this rejection is rendered moot.

The Examiner rejected claims 20-21 under 35 U.S.C. § 103(a) as being unpatentable over U.S. Patent No. 6,607,620 to Knickerbocker et al. in view of U.S. Patent No. 5,130,067 to Flaitz et al. as applied to claim 19, and further in view of JP 49-37921. Applicants traverse the rejection to the extent that it can be maintained.

Independent claim 20 and dependent claims 21, 70 and 71 relate to a method for reducing shrinkage during sintering low temperature co-fired ceramics. A layer to constrain shrinkage in the X-Y direction is positioned on a dielectric body to enclose an active area. The constraining layer is a low sintering temperature material comprising a strong auxiliary component to lower the sintering temperature of the layer. Bonding glass may be used to bond the constraining layer to the ceramic layer. Sintering involves bonding of adjacent surfaces of a mass of particles by heating below the fusion temperature. In sharp contrast, glass formation involves heating of glass-forming materials to produce a completely fused melt that is cooled to a non-crystalline molecular structure. Applicants enclose a page from A. Dodd and D. Murfin, "Dictionary of Ceramics", The Institute of Materials (1994); and two pages from W. Kingems, H. Bowen, and D. Uhlmann, Introduction to Ceramics, John Wiley & Sons (1991).

Knickerbocker et al. disclose a method for constraining shrinkage during processing of greensheets <u>prior</u> to sintering. Knickerbocker et al. apply a constraining frame that may be a glass ceramic material. Knickerbocker et al. fail to teach a constraining frame comprising a low sintering temperature material. As explained above, sintering involves forming an agglomeration of particles, whereas glass ceramic is a non-crystalline material. Applicants respectfully submit that glass ceramic does not suggest a mass of agglomerated particles. Further, the constraining frame is cut away before sintering the greensheets and therefore does not affect shrinkage during sintering (column 4 lines 36-53).

Flaitz et al. disclose a process for controlling Z-direction camber and X-Y bulge and distortion by applying pressure to the surface of a green ceramic structure during sintering. Flaitz et al. disclose several embodiments of their invention including the use of "contact sheets" that rely on friction to control shrinkage (beginning at column 8 line 65). The contact sheets must not fuse to the ceramic and are removed from the ceramic after sintering. There is no teaching or suggestion by Flaitz et al. of a process that controls shrinkage during firing of a monolithic structure that includes bonding glass applied between the contact sheet (constraining layer) and the ceramic layer (dielectric body). The combination Knickerbocker et al. and Flaitz et al. fail to teach or suggest a constraining layer (frame) comprising a low sintering material that is present during sintering; and fail to teach or suggest a strong auxiliary component to lower the sintering temperature of the constraining layer.

JP '921 is cited for teaching that glass can be strengthened by the presence of vanadium oxide. The Office Action concludes that it would be obvious to one of ordinary skill to similarly modify sintered powder. JP '921 also states that the presence of vanadium oxide provides glass having low expansibility, whereas the claimed sintered constraining layer reduces shrinkage of a ceramic substrate (paragraph [0043]). As explained by the "Dictionary of Ceramics" and the Introduction to Ceramics cited above, glass is a non-crystalline solid. It is usually formed from a melt. Because of a rapid cooling rate, glass has not enough time to form a long range ordered crystalline structure. Therefore, it is a non-crystalline solid. In sharp contrast, vanadium oxide in the present invention directly functions as a sintering aid to lower sintering temperature of refractory ceramics (page 12, paragraph [0037] and [0044]). Vanadium oxide is a crystalline material as opposed to non-crystalline glass. Vanadium oxide has a melting point of about 690 of the contrast of the present invention of the contrast of the contrast of the present invention of the contrast of the

C. After melting, it acts as a liquid phase to enable a liquid phase sintering. There is no reasonable expectation that a substance useful for modifying the physical properties of non-crystalline glass would provide a similar modifying effect on the physical properties of sintered particles. No art is cited that establishes that vanadium oxide performs an equivalent function in materials as physically and chemically diverse as glass and sintered powder. Factual findings, fully explained, to support the conclusions of the Office Action must be provided. See *In re Lee*, 61 USPQ2d 1430 (Fed. Cir. 2002).

Applicants respectfully submit that Knickerbocker et al. in view of Flaitz et al. and further in view of JP 49-37921 does not teach or suggest all of the elements of the invention as claimed, and request that the rejection on this ground be withdrawn.

The Examiner rejected claims 45-46 under Section 103(a) as being unpatentable over U.S. Patent No. 6,607,620 to Knickerbocker et al. in view of U.S. Patent No. 5,130,067 to Flaitz et al. as applied to claim 17, and further in view of U.S. Patent No. 6,488,795 to Sakai. Applicants traverse the rejection to the extent that it can be maintained. As these claims have been canceled, this rejection is rendered moot.

The Examiner rejected claims 47-49 under Section 103(a) as being unpatentable over U.S. Patent No. 6,607,620 to Knickerbocker et al. in view of U.S. Patent No. 5,130,067 to Flaitz et al. as applied to claim 17, and further in view of JP 2001-158670. Applicants traverse the rejection to the extent that it can be maintained. As these claims have been canceled, this rejection is rendered moot.

The Examiner rejected claims 67-68 under Section 103(a) as being unpatentable over U.S. Patent No. 6,607,620 to Knickerbocker et al. in view of U.S. Patent No. 5,130,067 to Flaitz et al. as applied to claim 19, and further in view of JP 58-115838 Abstract. Applicants traverse the rejection to the extent that it can be maintained. As these claims have been canceled, this rejection is rendered moot.

Claims 21, 70 and 71 depend directly or indirectly from claim 20. As explained above, claim 20 is now allowable over the combined art relied on in the Office Action. Because this

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amendment places the claims into condition for allowance, and raises no new issues, consideration and entry of this Amendment After Final is appropriate.

In view of the above amendments and remarks, Applicants respectfully request a Notice of Allowance. If the Examiner believes a telephone conference would advance the prosecution of this application, the Examiner is invited to telephone the undersigned at the below-listed telephone number.

23552

26 April 2006

Respectfully submitted,

MERCHANT & GOULD P.C. P.O. Box 2903 Minneapolis, Minnesota 55402-0903 (612) 332-5300

Brian H Batzli Reg. No. 32,960

## From A. Both and D. Murfin," Dictimary of Connics, The Institute of Historials Glaceranile, A lette that his been used

in USA for devittified glass products of the type exemplified by PYROCERAM Death Valley, California, (118 Boray and herate produced at the Geraley Mine, Geraley Barate, A sodium-calcium

Clarimeter, Sec INGERSOLL

Getting. The actual process of digging

Clemical Gorp., Los Angeles)

clay, by hand or by exervator; getting

and trånsporting form the successive

media har colours. (Allied Colloids, UK). Glas-Iok, Tradename. A glass-fibre joint Glascol, Tractename. Acrylic copolymer used as sealers for porous glazes and as for clay sewer pipes. (Pomona Pipe, GLARINGTER.

the Congo, and some other areas where

analysis is 65%, Al-O+ 33% H2O and

2% impurities, but many samples

contain clay. Gibbsite requires

Isterization has occurred. A typical

Aloy, 3H, O. Occurs in Dulch Guiana,

Gibbsite, Aluminium trihydrate,

states of winning.

rigid condition without drystallizing, and notes that glasses are typically hard and Glass. A solid with no long-range order in the arrangement of its aloms. ASTIM C162-92 defines a glass as an inorganic brittle, with a concholdal fracture. The word is used loosely as a synonym for product of fusion that has cooled to a

DONDED)

GLASSWARE (q.v.) The Annual Book of products. The names of many types of glass indicate their main constituents, ASTM standards Vol 15.02 lists over requirements for glass and glass fifty test methods and property calcination at a very high temperature to

gold, this is subsequently fired on at

ıbαıı 700⁼C.

made when W. H. Zachariasen (J. Amer. Chem. Noc., 54, 3841, 1932) deduced that silica being understood); for the system ni designating types of optical plass sec major advance in the understanding of are explicable if the interatomic forces the fundamental nature of a glass was the characteristic properties of glasses are essentially the same as in a crystal, e.g. soda-lime glass (the presence of OPTICAL GLASS CLASSIFICATION. A but if the three-dimensional atomic Gild. The painting of ponery with Equid climinate all the shrinkage that results FIFERNAL ENTANSION FACTORS FOR

of two londed rock which slide vertically determination of the initial and final set Cements and Monars, New York, 1804; The present form of apparatus consists needle for the final set is 1/2 in. (1 mm) Pricuent Treatise on Links, Hydrathic determination of initial set is 1/2 in. (2 dia. and weighs 1th (446g). Details are mm] dia. and weighs 1/2 lh (110g), the of portland cement (O. A. Gillmore. in a frame; the rud ('needle') for the Gilmore Needle. Apparatus for the givenlin ASTM-C206 (el. VICAT

old term for a tool used by carpenters to lumiture; cf. soxtusci. (From *Ginnet*, an chisa-ware, after 11 has been taken from Ginneter. Term in the N. Staffordshire the glost kiln, any adhering particles of polieries for a person who grinds from emove exerescences from wood.) refractory material from the kiln VITEDLE)

bonds with any other cations. The ions in oxygen atoms must be linked to only two pressure. This may be done by hand and a glass are thus divided into NGTWORK. piocesses: see blow-and-blow; piless only corners with each other; (3) some such cations and must not form further MODEFYING (q.v.). See also STRUCTON; Glassblowing, Shaping hot glass by air Glass-bonded Mico. See MICA (GLASSmay be an element of various forming mouth by a craftsman glassifiower, or VITROM, See also marine allasses, FORMING (Q.Y.) and NETWORK-AND-DLOW; PUFF-AND-BLOW. CHALCOGENIDE GLASSES.

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is subsequently devitrified in a controlled previously added in small amounts to the erystalline. The crux of the process is the can thus be varied, but 19 pically the ware conventional glassmaking processes, and and good thermal-shock resistance. Uses lemperature bearings, Unread-guides and glass batch, the nucleated article is then nucleated erystals can grow. Devittifiedmanner so that the linished product is glass products can be made in a wide heated to a temperature at which the range of compositions; the properties is impermeable and has high strength molten glass, is shaped by one of the material that, while in the form of a precipitation, during cooling of the domestic ovenware. (See also NEO-Glass-Ceramic. A type of ceramic shaped ware, of nucleating agents inchilde RAPONES (q.v.), high-CERAMIC GLASS).

has good resistance to chemical attack at netallic oxides, or salts, and low-melling A grade of vitroous enamelted ware that Glass-coated Steel or Glass-lined Steel high lemperatures and pressures; also Glass Enginel, A mixture of coloured known as Chassed Steet.

of cations that are surrounded by exygen

sample naist contain a high proportion

nejwajk, Zachanasen's rules for the formation of an exide glass are (1) the

network in a glass lacks lig symmetry

and periodicity of the crystalline.

these tetrahedra or triangles must share

retruhedra or by oxygen triangles; (2)

llux, e.g. kad horosiliente, used for the decoration or labelling of glass-ware; the spinning, or by the telion of a blast of air furnace to form tilaments. Nowadays the Owens stillen-phowits a profess (p.s.) Glass Fibre. Filamentous glass made by Glass Live. A large unbroken blister on glass filtres are known as 'Sik', a fleece because free of surface flaws. There are over 20 British Standards for glass fibre products with glass libre reinforcement. making libres was produced as marbles bush, at winding speeds of 50 m/s. The or steam to produce Suph Films. The bornded ingether form a Sumul. Long direct-unch method is used. Fibres are blaments are a few pm diameter, and drawn directly from holes in an alloy unprocessed filaments are known as Glass Frost, Very thin this that bus been crushed for use as a decorative material is 'Mat', Formerly glass for mechanical drawing or centrifugal which were remelted in an electric Basic Fibre; a miniber of filtiments much stronger than ordinary glass like mass of libres is Wood; felly is used to make glass word Cilise products and plastics and ecotoni Glass-energysulated Hilling, Sec enamel is lited on at a red heat. ENCARSULATED HIPING vitreous enamelware. malerial (cf. Trysia ).

encapsulation in plays. An Americanism and polymers and copolymers of acryb between aluministicate glass panders. Government Chemist. (J. Appl. Chem. denial cements, made by the reaction Glass-tanonner Cements, Translucent acid. Developed by A.D. Wilson and semiconductor devices by complete B.E. Kent at the Laboratory of the Biotechnol 21, (11), 315, 1971). Glassivation, Passivation of specification, see B.S. 6039.

Gilard and Duhrul Fuctors, See

from the loss of water.

NTRODUCTION TO CERAMICS

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Sificates bare structures in which SiO," letrakedra share vertices to form chains, rags, sheets, etc. In phosphates 4PO.") and solfates (SO.") similar tetralisers are found, but they are always included. Yet AIPO, has a structure

The alkabra earth oxides. MgO, SrO. BaO, all base the rock sall structure. The which curresponds to that of quarts, SiO, Ê

Mro (each salt uncerture) and Liso familiamine structure) both are based on cubic close parked oxygen, with cations ocrupying interstices in the saray. Yel psedominant point Gelect in MgO is of the Schottky type: Mat in L.,O Is of the hardness and melting points of the compounds decreases in the order given. Frenkel type. 3

The garnets Mg.Aki/SiO, h and Fe, Al, (SiO,), are is unvertibous in a manner similar to My,510, or Fe,5.0, isomorphous with Ca,510. Give an explanation for this based on Fessio, and Alg.SiO. They are not isomorphous with CasAlg(SiO.),, and neither is imis size and coordinating numbers. On the basis of your theory predict a mineral not mentioned in this question that will be isomorphous with MaSiCa, one for Cz.SiO., one for Ma.A.J.SiO.h. and one for Ca,A.J.SiO.h. Ξ

erystalized from a glass rack. The X-ray diffraction pattern showed they were a A centain enumers was asked to identify some place-shaped exposuls that had single phase fonly one kind of crystal structure), but the chanical analysis indicated a called you in as a consultant, would you be able to show him this is related to complicated formula KF-AIF-BaO-MgO-AI,O,-SNgSiO, on an empirical basis. If he (postassium mica) and to the tale or pyrophillite erystals? Show what substitutions in tale or pyrophillite that have been made to produce this crystal. Ξ.

Introduction to Cermanies," John Wiley & Sous From Witingery, HiBowen, D. Uhlinam," က

Structure Glasses

Even though a majority of natural and manufactured solids are crystalline are of great importance for both traditional and newly deformulation of glasses, glazes, and enamels. Solid plasses, of which the silicates are the technologically most important group, usually have a more complex structure than the liquids from which they are derived, and more newly developed class of materials is thin films deposited us materials the short-range order is preserved in the immediate vicinity of ized by diversity among different systems and by difficulty in precise crystalline in nature, as discussed in Chapter 2, materials which are not veloped ceramics. One important chass is the liquid silicates, the properlies of which are an essential part of the ceranist's knowledge in the recent studies indicate a complexity which is still not well understoad, although the broad structural characteristics seem reasonably clear. A בי ביים חשיים בינות בינות בינות מות שבים בינות ב known as to structural details. In each of these classes of ceramic any selected atom, that is, the first coordination ring; the longer-range order characteristic of the ideal crystal is dissipated in a way characterdescription.

. We focus our aftention on glasses, which pre, by far, the most insportant. group of inorganic noncentralinessolids. The structure of glasses may be considered on three scales: (1) the scale of 2 to 10 Å, or that of livent atomic arrangements; (2) the scale of 30 to a few thausand angstroms, or that of submicrostructure; and (3) the scale of nicrons to millimeters or more, or that of micrustructure and macrostructure. In this chapter we consider the atomic structure and the submicrostructure of glasses; the consideration of microstructural features is deferred to Chapter 11. 5

## STRUCTURE OF GLASSES

for the structure to relax is increased, the supercooled liquid persists to n TYSIBILINE solid. If slower cooling rates are used so that the time available ituclure does not relax at the cooling rate used. The expansion wedlitient for the glassy state is usually about the same as that for the lower temperature, and a higher-density glass results, Similarly, by benting the glassy material in the annealing range, in which slow returntion can occur, the glass structure in time approaches an equilibrium density corresponding to the supercooled liquid at this temperature. ninginguithing

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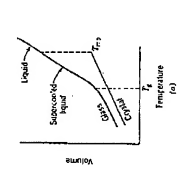
A concept useful in discussing the properties of glasses is the glass intersection between the curve for the glassy state and that for the supercoaled Isquid (Fig. 3.1), Different cooling rates, corresponding to transition temperature T, which corresponds to the temperature of the different relaxation times, give rise to a different configuration in the glassy state equivalent to different points along the curve for the supercooled liquid. In the transition range the time for structural rearrangements is sinilar in magnitude to that of experimental observations. Consequently the configuration of the glass in this temperature range changes slowly with time toward the equilibrium structure. At somewhat higher temperatures the structure corresponding to equilibrium at any temperature is achieved very rapidly. At substantially lower temperatures the configuration of the glass remains sensibly stable over long periods of time.

In discussing the structural characteristics of glasses, reference is often made to the structure of a particular glassy material. It should be nowed, however, that any determination of glass structure is only meaningful has increased to a sufficiently high value, typically about 10" to 10" P, so within limits seen from the volume-temperature relations shown in Fig. 3.1. As the liquid is cooled from a high temperature without crystallizing. a region of lemperature is reached in which a bend appears in the volume-temperature relation. In this region, the viscosity of the material that the sample exhibits solidlike behavior. As shown in Fig. 3.1b, the glass transition temperature increases with increasing cooling rate, as do the specific volumes of the glasses which are formed. In the case shown, the specific volume of the glass at temperature  $Y_a$  can be  $V_1$  or  $V_2$  or  $V_4$ depending on which of the three cooling rates was used in forming the glass. The maximum difference in specific volume obtainable with variations in the cooling rate is typically in the range of a few percent; only within this range can one speak of the structure of a glass without carefully specifying its mode of formation,

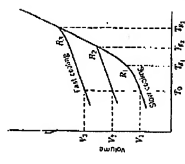
the liquid state, and their structure may differ significantly from glasses Noncrystalline solids can be formed in other ways besides cooling from ormed by the cooling of liquids. Among these alternative methods, the

seen by a plot of the specific volume of the crystal, liquid, and glass as a function of temperature (Fig. 3.1). On cooling the liquid, there is a decreases at about the same rate as above the melting point until there is a decrease in the expansion coefficient at a range of temperature called the structure of glasses can be clearly distinguished from that of liquids, since lizes. However, if no crystallization occurs, the volume of the liquid transformation range. Below this temperature range the glass glass structure is effectively independent of temperature. This can best be discontinuous change in volume at the melting point if the liquid crystalby solidification from the INTRODUCTION TO CERAMICS ormed Glasses are usually Glass Formation

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crystal; (b) glastes formed at different cooling Fig. 3.1. Schematic specific volume-tempenaliquid, glass, and ture relations. (a) Relations for rates R, < R, < R,

Temperature

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